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(54) Bleach manganese catalyst and its use.

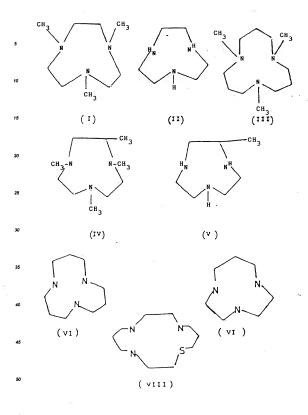
(iii) Novel bleach and oxidation catalysts are reported as well as a bleaching-detergent composition employing these catalysts and a method for removing stains from substrates. The catalysts are manoanese complexes of the formula (A):

[LMn* (OR)] Y, wherein mn is manganese in the +4 oxidation state;

R is a Cri-Coo radical selected from alkyl, cycloalkyl, aryl, benzyl and radical combinations thereof; at least two R radicals may also be connected to one another so as to form a bridging unit between two oxygens that coordinate with the manganese;

L is a ligand selected from a C₂-C₆₀ radical having at least 3 nitrogen atoms coordinating with the manganese; and

Y is an oxidatively-stable counterion.



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HN(CH₂CH₂NH₂)₂ XI CH₃N(CH₂CH₂N(CH₃)₂)₂ XII CH₃C(CH₃CH₃N(CH₃)₂)₃ XIII

The most preferred ligands are ligands with (I) to (V) being particularly preferred.

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Ligend (I) is 1,4,7-trimethyl-1,4,7-triazacyclononane, coded as Me₂-TACN; ligand (II) is 1,4,7-triazacyclononane, coded as TACN; ligend (III) is 1,5-trimethyl-1,5,9-triazacyclododecane, coded as Me₂-TACN; ligend (IV) is 2-methyl-1,4,7-trimethyl-1,4,7-triazacyclononane, coded as Me/Me₂-TACN; end ligend (V) is 2-methyl-1,4,7-triazacyclononane, coded as Me/TACN.

Any of these complexes, are useful catalysts for the bleech ectivation of peroxy compounds over e wide class of stains et lower temperatures in e much more effective wey than the Mr-based catalysts of the ert hitherto known. Furthermore, these catalysts exhibit a high stability egainst hydrolysis and oxidation, even in the presence of oxidants such as hypochlorite.

Another edventage is that, in many respects, the instant cetalysts are better than eny other Mn-complexes proposed in the ert. They are not only effective in enhancing the bleaching ection of hydrogen peroxide bleaching agents but less of orgenic and inorgenic peroxyacid compounds.

A surprising feature of the bleach systems according to the invention is that they ere effective on e wide range of stains including both hydrophilic and hydrophobic steins. This is in contrast with all previously proposed Mn-besed catalysts, which are only effective on hydrophilic stains.

A further surprising feature is that they ere competible with detergent enzymes, such as proteases, cellulases, lipases, amylases, oxidases etc.

The invention, therefore, provides a bleeching or cleaning processs employing a bleeching egent selected from the group of peroxy compound bleaches including hydrogen peroxide, hydrogen peroxide liberating or generating compounds, peroxyecids end their salts, end peroxyecid bleach precursors and mixtures thereof, which process is characterised in that the bleaching agent is activated by e catalytic amount of e Mn-complex of general formula (A) as defined hereinbefore.

The catalytic component is a novel feature of the invention. The effective level of the Mr-complex catalyst, expressed in terms of perts per million (ppm) of manganese in the aqueous bleaching solution, will normelly range from 0.001 ppm to 100 ppm, preferably from 0.01 ppm to 20 ppm, most preferably from 0.1 ppm to 10 ppm. Higher levels may be desired and applied in industrial bleaching processes, such as textile and paper pulp-bleeching. The lower range levels ere primarily destined end preferably used in domestic leundry end dishwashing operations. Levels of peroxy compound in the aqueous solution will range from 0.01 ppm to 250 ppm, portinally from 0.5 ppm to 100 ppm.

The improved bleeching composition of the invention has particular application in detargent formulations to form a new and improved detergent bleach composition within the purview of the invention, comprising the peroxy compound bleach, the eforesaid Mn-complex catalyst, a surface-active material, and usually also detergency builders and other known ingredients of such formulations, es well as in the industrial bleaching of yerns, laxtiles, paper, woodpulp and the like.

The Mr-complex catalyst will be present in the detergent formulations in amounts so es to provide the required level in the wash liquor. When the dosage of the detergent bleech composition is relatively low, e.g., about 1 and 2 g/l by consumers in Japan and the USA, respectively, the Mn content in the formulation is 0.0025 to 0.5%, preferably 0.005 to 0.25%. At higher product dosage as used e.g. by European consumers, the Mn content in the formulation is 0.0005 to 0.1%, preferably from 0.001 to 0.05%.

Compositions comprising e peroxy compound bleech end the eforesaid bleech catalyst ere effective over a wide pH range of between 7 end 13, with optimal pH range lying between 8 and 11.

The peroxy compound bleaches which can be utilized in the present invention include hydrogen peroxide, hydrogen peroxide-liberating compounds, hydrogen peroxide-generating systems, peroxyacids and their salts,

and peroxyacid bleach precursor systems, and mixtures thereof.

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Hydrogen peroxide sources are well known in the art. They include the alkali metal peroxides, organic peroxide bleaching compounds such as urea peroxide, and inorganic perate bleaching compounds, such as the alkali metal perborates, percarbonates, perphosphates and persulphates. Mixtures of two or more of such compounds may also be suitable. Particularly preferred are sodium percorate and odium perborate and objectively, sodium perborate monohydrate. Sodium perborate monohydrate is preferred to tetrahydrate because of its excellent storage stability while also dissolving very quickly in aqueous bleaching solutions. Sodium percorate monohydrate is perferred to revinormental reasons. These bleaching compounds may be utilised alone or in conjunction with a peroxyacid bleach precursor. Use of precursors may be of advantage for improving the overall whiteness appearance of white fabrics as well as for hygiene purposes.

Peroxyacid bleach precursors are known and amply described in the literature, such as in the GB Patents 888, 884,798; 97,356; 1,003,310 and 1,519,351; German Patent 3,337,321; EP-A-0185522, EP-A-0174132; EP-A-0120591; and U.S. Patents 1,246,339; 3,322,882; 4,128,494; 4,412,934 and 4,575,393.

Another useful class of peroxyacid bleach precursors is that of the quaternary ammonium substituted peroxydid precursors as disclosed in U.S. Patenta 4,76,1015 and 4,397,757. In EP-A-284292, EP-A-331229, EP-A-303520, EP-A-453398 and EP-A-464880. Examples of peroxyacid bleach precursors of this class are:

2-(N,N,N-trimethyl ammonium) ethyl-4-sulphophenyl carbonate - (CSPC);

N-octyl,N,N-dimethyl-N₁₀-carbophenoxy decyl ammonium chloride - (ODC);

3-(N,N,N-trimethyl ammonium) propyl sodium-4-sulphophenyl carboxylate; and N,N,N-trimethyl ammonium toluyloxy benzene sulphonate.

Of the above classes of bleach precursors, the preferred classes are the esters, including acyl phenol sulphonates and acyl alkyl phenol sulphonates; acyl-amides; and the quaternary ammonium substituted peroxvacid precursors.

Highly preferred activators include sodium-4-benzoyloxy benzene sulphonate; N.N.N'.N'-tetraacetyl ethylene diamine; sodium-1-methyl-2-benzoyloxy benzene-4-sulphonate; sodium-4-methyl-3-benzoyloxy benzoate; trimethyl ammonium toluyloxy benzene sulphonate; sodium nonanoyloxybenzene sulphonate; sodium 3,5,5-trimethyl hexanoyloxybenzene sulphonate; glucose pentaacetate and tetraacetyl xylose.

Organic peroxyacids are also suitable as the peroxy compound. Such materials normally have a general formula:

wherein R is an alkylene or substituted alkylene group containing from 1 to about 22 carbon atoms or a phenylene or substituted phenylene group, and Y is hydrogen, halogen, alkyl, aryl or

The organic peroxy acids usable in the present invention can contain either one or two peroxy groups and can be either aliphatic or aromatic. When the organic peroxy acid is aliphatic, the unsubstituted acid has the general formula:

where Y can be, for example, H, CH₃, CH₂CI, COOH, or COOOH; and n Is an integer from 1 to 20.

When the organic peroxy acid is aromatic, the unsubstituted acid has the general formula:

wherein Y is hydrogen, alkyl, alkylhalogen, halogen, or COOH or COOOH.

Typical monoperoxy acids useful herein include alkyl peroxy acids and aryl peroxy acids such as:

- i) peroxybenzoic acid end ring-substituted peroxybenzoic ecids, e.g. peroxy-α-naphthoic ecid;
- ii) aliphatic, substituted aliphatic and arylalkyl monoperoxy acids, e.g. peroxylauric acid, peroxystearic acid, and N,N-phthaloylaminoperoxyceproic acid (PAP).

Typical diperoxy acids useful herein include elkyl diperoxy ecids end aryldiperoxy acids, such as:
iii) 1.12-diperoxydodecanedioic acid;

iv) 1.9-diperoxyazelaic acid;

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- v) diperoxybrassilic ecid; diperoxysebasic acid end diperoxyisophthalic ecid;
- vi) 2-decyldiperoxybutane-1,4-dioic ecid;
- vii) 4.4'-sulfonylbisperoxybenzoic ecid.
- An inorganic peroxyacid salt useble herein is, for example, potassium monopersulphate.

A detergent bleach composition of the invention can be formuleted by combining effective emounts of the components. The term "effective amounts" es used herein means that the ingredients are present in quantities such that each of them is operative for its intended purpose when the resulting mixture is combined with water to form an aqueous medium which can be used to wash end deen dothes, fabrics end other articles.

In particular, the detergent bleach composition can be formuleted to contain, for example, from ebout 2% to 30% by weight, preferably from 5 to 25% by weight, of e peroxide compound.

Peroxyacids may be utilized in somewhat lower amounts, for example from 1% to about 15% by weight, preferably from 2% to 10% by weight.

Peroxyacid precursors mey be utilized in combination with a peroxide compound in approximately the same level as peroxyacids, i.e. 1% to 15%, preferably from 2% to 10% by weight.

The manganese complex catalyst will be present in such formuletions in emounts so as to provide the required level of Mn in the wash liquor. Normally, an emount of menganese complex catalyst is incorporated in the formulation which corresponds to a Mn content of from 0.0005% to about 0.5% by weight, preferably 0.001% to 0.25% by weight.

The bleach catalyst of the invention is compatible with substantially any known and common surface-active agents and detergency builder meteriels.

The surface-active material may be naturally derived, such es soap, or a synthetic material selected from anionic, nonionic, amphoteric, zwitterionic and cationic actives and mutures thereof. Many suitable actives ere commercially available and are described in the literature, for example in "Surface Active Agents and Detergents", Volumes I end II, by Schwartz, Perry and Berch. The totel level of the surfece-active material may range up to 50% by weight, preferebly being from about 1% to 40% by weight of the composition, most preferably 4 to 25%.

Synthetic anionic surface-ectives ere usually watersoluble elkali metal salts of organic sulphates and sulphotates having allkyl groups containing from about 8 to about 22 carbon etoms, the term alkyl being used to include the alkyl portion of higher anyl groups.

Examples of suitable synthetic enionic detergent compounds ere sodium end armonium alkyl sulphates, especially those obtained by sulphating higher $(C_p\cdot C_{10})$ alcohols produced, for example, from tallow or occonut oil, sodium and armonium alkyl $(C_p\cdot C_{20})$ because sulphonates, particulerly sodium linear secondary elkyl $(C_p\cdot C_{10})$ because sulphonates; sodium alkyl glyceryl ether sulphates, especially those esters of the higher alcohols derived from tallow or occonut oil end synthetic alcohols derived from tallow or occonut oil afty acid monoglyceride sulphates and sulphonates; sodium and armonium selts of sulphuric acid esters of higher $(C_p\cdot C_{10})$ fatty alcohol alkylene oxide, particularly ethylene oxide, reaction products; the reaction products of talty acids such as coconut fatty acids esterified with isethionic acid and neutralised with sodium hydroxide; sodium and armonium salts of fatty acid amides of methyl tautine; alkene monosulphonetes such as those derived by reacting paraffins with SO_2 and CI_2 and then hydrolysing with a base to produce C_1 and such acids and C_2 and C_3 and C_4 and then hydrolysing with a base to produce a random sulphonate; sodium and armonium C_1 C_2 allows sulfosuccinates; C_3 elphaolefins, with SO_3 and then neutrelising and hydrolysing the reaction product. The preferred anionic detergent compounds are sodium $(C_1 - C_{10})$ elkylbenzenes sulphonates, sodium $(C_2 - C_{10})$ elkylbenzenes sulphonates.

Examples of suitable nonionic surfece-active compounds which mey be used include, in particuler, the reaction products of alkylene oxides, usually ethylene oxide, with alkyl (C₈-C₂₉) phenols, generally 5-25 EO, i.e. 5-25 units of ethylene oxides per molecule; the condensation products of eliphetic (C₉-C₁₉) primery or secondary lineer or branched alcohols with ethylene oxide, generally 3-30 EO, and products made by condensation of ethylene coxide units oxide with effect oxide with experiment oxide alkylene diamine. Other so-called nonionic surface-actives include alkyl polyglycosides, long chain tertiary emine oxides, long chein tertiery phosphine oxides end delikyl subhoxides.

Amounts of amphoteric or zwittenonic surface-ective compounds cen elso be used in the compositions of the invention but this is not normally desired owing to their relatively high cost. If any emphoteric or zwittenonic

datargant compounds are used, it is generally in small amounts in compositions based on the much more commonty used synthetic anionic and nonionic activas.

As stated above, soaps may also be incorporated in the compositions of the invention, preferably at a level of less than 25% by weight. Thay are particularly usaful at low levels in binary (soap/anionic) or ternary mixtures together with nonionic or mixed synthatic anionic and nonionic compounds. Soaps which are used, are prafarably the sodium, or, less desirably, potassium salts of saturated or unsaturated C10-C24, fatty acids or mixtures thereof. The amount of such soaps can be varied between about 0.5% and about 25% by weight, with lower amounts of about 0.5% to about 5% being generally sufficient for lather control. Amounts of soap between about 2% and about 20%, especially between about 5% and about 10%, are used to give a banaficial affect on detergency. This is particularly valuable in compositions used in hard water when the soap acts as a supplamantary buildar.

The detargent compositions of the invention will normally also contain a datargency builder. Builder materials may be selected from 1) calcium sequestrant materials, 2) pracipitating materials, 3) calcium ion-exchange materials and 4) mixtures thereof.

Examplas of calcium sequestrant buildar materials include alkali matal polyphosphates, such as sodium tripolyphosphata; nitrilotriacetic acid and its water-soluble salts; the alkali metal salts of ather polycarboxylates, such as carboxymethyloxy succinic acid, oxydisuccinic acid, mellitic acid; ethylene diamine tetraacetic acid; benzene polycarboxylic acids; citric acid; and polyacetal carboxylates as disclosed in U.S Patant No.s 4,144,226 and 4,146.495.

Examples of precipitating builder materials include sodium orthophosphate, sodium carbonate and sodium carbonate/ calcite.

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Examples of calcium ion-exchange builder materials include the various types of water-insoluble crystalline or amorphous aluminosilicates, of which zeolites are the best known representatives.

In particular, the compositions of tha invention may contain any one of tha organic or inorganic builder materials, such as sodium or potassium tripolyphosphate, sodium or potassium pyrophosphate, sodium or potassium orthophosphata, sodium carbonate or sodium carbonate/ calcite mixtures, the sodium salt of nitrilotriacatic acid, sodium citrata, carboxymethyl malonata, carboxymethyloxy succinate and the water-insoluble crystalline or amorphous aluminosilicate buildar materials, or mixturas theraof.

These builder materials may be present at a level of, for example, from 5 to 80% by weight, preferably from 10 to 60% by weight.

Apart from the components already mentioned, the detargent compositions of the invention can contain any of the conventional additivas in the amounts in which such materials are normally amployed in fabric washing datergant compositions. Examplas of thase additivas includa lathar boostars, such as alkanolamidas, particularly the monoethanol amides darived from palmkarnal fatty acids and coconut fatty acids, lather daprassants, such as alkyl phosphatas and silicones, anit-radaposition agents, such as sodium carboxymathyl cellulosa and alkyl or substituted alkyl celluose ethers, other stabllizers, such as athylana diamina tatraacatic acid and the phosphonic acid derivatives (i.a. Dequest^R types), fabric softening agants, inorganic salts, such as sodium sulphate, and usually present in very small amounts, fluorescent agents, perfumas, enzymes, such as proteases, cellulasas, lipasas, amylases and oxidasas, germicidas and colorants.

Another optional but highly desirable additive ingredient with multifunctional characteristics in datergent compositions is from 0.1% to about 3% by weight of a polymeric material having a molacular weight of from 1,000 to 2,000,000 and which can be a homo- or copolymer of acrylic acid, maleic acid, or salt or anhydride thereof, vinyl pyrrolidona, methyl- or ethylvinyl athers, and other polymerisable vinyl monomers. Preferred examplas of such polymaric matarials are polyacrylic acid or polyacrylate; polymalaic acid/acrylic acid copolymar; 70:30 acrylic acid/hydroxyethyl maleate copolymar, 1:1 styrena/maleic acid copolymar, isobutylena/maleic acid and diisobutylene/malaic acid copolymers; mathyl- and ethylvinyl ather/maleic acid copolymars; ethylene/malaic acid copolymer; polyvinyl pyrrolidone; and vinyl pyrrolidone/maleic acid copolymar.

Detergent bleach compositions of the invantion, when formulated as free-flowing particlas, a.g. in powderad or granulated form, can be produced by any of the conventional techniques employed in the manufacture of detargent compositions, for instanca by slurry-making, followed by spray-drying to form a detergent base powder to which the heat-sansitive ingredients including tha peroxy compound blaach and optionally some other ingredients as desired, and the bleach catalyst, can be added as dry substances.

It will be appreciated, however, that the datargent base powder compositions to which the bleach catalyst is added can itsalf be made in a variety of other ways, such as the so-called part-part processing, non-tower routa processing, dry-mixing, agglomeration, granulation, axtrusion, compacting and densifying processas atc., such ways being wall known to those skilled in the art and not forming the assential part of the present invention.

Advantageously, it is useful to separate the manganese complex from the peroxygen compound and other

components of the detergent-bese powder. This may be eccomplished through granulation or noodling (through en extruder) the managanese complex within e protective matrix shell. Sodium sulfate and other inorganic or organic substances elong with binders may be useful in forming the mangenese complex protective perticles.

Alternetively, the bleach catalyst can be edded seperately to a wash/bleach weter containing the peroxy compound bleeching agent.

In the case, the bleech catalyst is present as a detergent additive product. Such additive products ere intended to supplement or boast the performance of conventional detergent compositions end mey contain eny of the components of such compositions, although they will not comprise ell of the components as present in a fully formuleted detergent composition. Additive products in accordance with this espect of the invention will normally be added to en aqueous illegor containing a source of (alkeline) hydrogen peroxide, elthough in certain circumstances the additive product may be used as separate treatment in e prewash or in the rinse.

Additive products in eccordance with this espect of the invention mey comprise the compound alone or preferably, in combination with a carrier, such es a compatible equeous or nonaqueous liquid medium or e particulate substrate or effective nonparticulate substrate.

The following exemples will more fully illustrate the embodiments of this invention. All parts, percentages and proportions referred to herein end in the eppended claims ere by weight unless otherwise indicated.

EXAMPLES

Example 1

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Preparation of Mn(IV)Me₃TACN(OMe)₃(PF₆)

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Menganese (II) chloride was obtained from Strem Chemicals, Inc. Sodium hexefluorophosphate was obtained from Aldrich Chemical Company, Sodium methoxide was prepared from sodium hydride (Aldrich) end methanol. N,N',N'-trimethy1, 1,4,7-triazacyclononene (Me₃TACN) was prepered eccording to the literature procedure from Atkins, et el. 'Orgenic Synthesis', 58, 86, 1978.

A degessed (duel-manifold vacuum line technique) solution of 2.5 g Me₃TACN in 40 ml methanol wes chilled in an iocseth bath. To this solution was edded 1.84 g manganese chioride dissolved in 10 ml methenol. The solution turned light brown. Subsequently, 20 ml of 1M NeOMe was edded ell at once whereupon the solution turned darker brown. After stirring for about 10 minutes, 5.0 g NaPF₆ (in 20 ml methenol) was edded, and the solution was exposed to eir. About 2-30 minutes leter, a further emount of 5.0 g NaPF₆ (in 20 ml methanol) was added, and the solution was ellowed to warm to room temperature. The solution was then filtered and cooled to 0°C to promote crystallisation. After three days, derk brown crystals formed which were filtered. Yield 0.5g (16% - besed on NaOMe). More crystals were obtained from later crystallisetions, but often these conteined co-crystallised est.

Elementel enalysis: Calculated for C₁₂H₃₀F₆MnN₃O₃P: C, 31.03; H, 6.47; Mn, 11.85; N, 9.05; P, 6.68. Found: C, 29.28; H, 5.68; Mn, 10.60; N, 9.20; P, 6.86. Mass spectral enelysis: m/e 319.

UV enalysis (CH₃CN): lambda max (nm) (E, L mol⁻¹ cm⁻¹) 326 (11,200), 287 (10,900), 228 (9800). Melting range: 160-165°C.

EXAMPLE 2

The bleach activating ability of Mn(IV)(Me₃TACN)(OCH₃)₃(PF₆) was demonstrated on Regu (i.e. e tomatobased product) steined end tea stained (BC-1) cotton cloths at verious active levels and various active oxygen levels.

Bleaching Terg-O-Tometer experiments were performed et 40°C using the recommended dose of detergent powder (1.55g/L of zeolite containing bese powder) in deionised, distilled weter, for e 15-minute wash with 2 or 4 stained oldshe per one litter pot. The bleeching catalyst (Mn catalyst) was edded se s 5% Ne₂SO₄ powder blend. The performance of the monomer, Mn(IV)(Me₂TACN)(OCH₃)₃(PF₆) was compered to the performence of enother manganese bleach catalyst, the dinuclear mengenese complex. Mn(IV)(Me, ACN)(LON)(LPO)(E), HyO. Results are listed under Table I.

Bleeching values are reported as changes in reflectance (R) or (B) units (LAB scale) as measured using a Colorgerd/System/05 Reflectometer. Since peroxide-only controls were elso conducted on the same type stains, bleeching was actuelly reported as $\Delta\Delta B$ and $\Delta\Delta B$ bit here colocaleted as $\Delta\Delta B$ = ΔR (wesh) - ΔR (blenk) and $\Delta\Delta B$ = - $\langle \Delta B |$ wash)- ΔR (blank) end, therefore, the higher the number, the better the performance.

Table 1

Ragu Bleading Studies (Mn catalysts)		
	Mn (IV) (Me ₃ TACN) (OCH ₃) ₃	Mn (IV) ₂ (Me ₃ TACN) ₂ (μ-O) ₃
	ΔΔR	ΔΔR
48 ppm [0]	9.7	0.6
24 ppm [0]	9.1	0.6
12 ppm [0]	6.7	1.2
6 ppm [0]	7.2	0.8

Conditions: 4 ppm Mn complex, 40°C, 15 minute wash, pH=10, 1.55g/l Zeolite Base, [0] = active oxygen added as sodium perborate.

These results indicate that the Mn monomer, Mn(IV)(Me₃TACN) (OCH₃)₃(PF₆), activates hydrogen peroxto remove the hydrophobic Regus tain. It performed much better than the corresponding dinuclear Mn catalyst at the same level of actives on this "0!ly" stain.

Example 3

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This Example presents the results of bleaching studies with Mn(IV)(Me₃TACN)(OCH₃)₃(PF₆) on tea stained cloth. Test values are reported in Table II.

Table II

	Table II	
Tea Stain (BC-1) E	Bleaching Studies (Mn Catalysts)	
	Mn (IV) (Me ₃ TACN) (OCH ₃) ₃	Mn (IV) ₂ (Me ₃ TACN) ₂ (μ-O) ₃
	ΔΔR	ΔΔR .
48 ppm [O]	21.2	18.5
24 ppm [O]	16.0	12.1
12 ppm [O]	11.7	8.4
6 ppm [O]	7.1	4.1

40 Conditons: 4 ppm Mn complex, 40°C, 15 minute wash, pH=10, 1.55g/l zeolite base powder, [O] = active oxygen added as sodium perborate.

These results again Indicate that the Mn monomer is a very efficient bleach and again is somewhat better at removing the tea stain than the corresponding dinuclear Mn bleach catalyst. Comparing the performance at various active oxygen levels also indicates that the monomer gives comparable bleaching of the tea stain at 30-50% lower active oxygen levels than the manganese dimer.

Example 4

This Example details results similar to that of Example 3 but evaluates the effectiveness of extremely low levels of catalysts. Test results are reported under Table III.

Table III

Tea Stain (BC-1) Bleaching Studies (Mn Catalysts)			
	Mn (IV) (Me ₃ TACN) (OCH ₃) ₃	Mn (IV) ₂ (Me ₃ TACN) ₂ (μ-O) ₃	
	ΔΔR	ΔΔR	
4 ppm Mn complex	21.1	18.5	
2 ppm Mn complex	16.0	12.1	
1 ppm Mn complex	12.1	10.0	

Conditions: 40°C, 15 minute wash, pH=10, 48 ppm active oxygen added as sodium perborate, 1.55g/l zeolite base powder.

These results show that at even lower catalyst levels sufficient bleaching is occurring and again is comparable to or somewhat better than the corresponding dinuclear manganese bleach catalyst.

Example 5

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This Example illustrates the effectiveness of Mn(IV)(Me₃TACN)(OCH₃)₃(PF_e) with respect to a variety of different bleachable stains. Test results are reported in Table IV.

Table IV

Staining Bleaching Studies	with Mn(IV) (Me ₃ TACN) ((OCH ₃) ₃	
	Perborate-only	Mn(IV) (Me ₃ TACN) (OCH ₃) ₃	
	ΔR(-ΔΒ)	ΔR(-ΔΒ)	
Tea Stain (BC-1)	-0.5	21.7	
Ragu	5.5	15.2	
Red Wine (EMPA-114)	13.2	35.1	

Conditions:

4 ppm Mn complex, 40°C, 15 minute wash, pH=10, 1.55g/l zeolite base powder, 48 ppm active oxygen added as sodium perborate.

The results in Table IV indicate that the Mn monomer, Mn(IV)(Me_TACN)(OCH₃)₃(PF₆), is effective on a variety of very different bleachable stains. The catalyst is effective on both hydrophobic and hydrophilic stains and performs at very low levels to clean and bleach very difficult stains.

Example 6

This Example illustrates the effectiveness of Mn(IV)(Me₃TACN)(OCH₃)₃(PF₆) as it performs in conjunction with different peracids. Test results are reported under Table V.

Table V

Catalysed Peracid Bleaching Studies with Mn(IV) (Me ₃ TACN) (OCH ₃) ₃		
	Without Catalyst	With Catalyst
	ΔΔR	ΔΔR
Peracetic Acid	9.7	26.0
Sodium Monopersulfate	29.8	37.6

Conditions: 4 ppm Mn complex, 40°C, 15 minute wash, pH=10, 1.55g/l zeolita basa powdar, 8 x 10⁻³M paracid, Taa stainad cloths (BC-1).

Thasa rasults show that Mn(IV)(Ma₂TACN)(OCH₃)₃(PF₆) affactivaly catalysas tha bleaching of paracids as well as hydrogen peroxide at very low lavals of catalyst (4 ppm) on tea stained doth. The activated paracetic acid performed almost three times as affectively as the paracid alona and the activated monopersulfate removed all of the tea stain from the doth leaving the initially brown doth totally white.

The foregoing description and examples illustrate selected embodiments of the present invention. In light theraof, various modifications will be suggested to one skilled in the art, all of which are within the spirit and purylew of this invantion.

Claims

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1. A catalyst having the formula (A):

[LMn^{[V} (OR)₃] Y, (A)

wherain Mn is manganasa in tha +4 oxidation stata;

R is a C_1 - C_{20} radical selected from alkyl, cycloalkyl, aryl, benzyl and radical combinations thereof, at least two R radicals may also be connected to one another so as to form a bridging unit between two oxygans that coordinate with the manganasy.

L is a ligand salactad from a C_3 - C_{60} radical having at least 3 nitrogen atoms coordinating with the manganesa; and

Y is an oxidatively-stable counterion.

- A catalyst according to claim 1 wherain tha ligand L is selacted from 1,4,7-trimathyl-1,4,7-triazacyclonoriana, and 2-methyl-1,4,7-trimathyl-1,4,7-triazacyclononane.
- A catalyst according to claim 1 which is Mn(IV) (Ma₃TACN) (OCH₃)₃ (PF₆).
- A bleaching composition comprising:

i) from about 1 to about 60% by weight of a paroxy compound; and

 ii) a manganesa complax present in an affective amount for catalysing bleaching activity of the peroxy compound and having the structure:

[LMn' (OR)3] Y, (A)

wherein Mn is manganesa in the +4 oxidation stata; R is a C_1 - C_{20} radical selected from alkyl, cycloalkyl, aryl, benzyl and radical combinations thereof;

at least two R radicals may also be connected to one another so as to form a bridging unit between two oxygens that coordinate with the manganesa;

L is a ligand selected from a C₃-C₈₀ radical having at least 3 nitrogan atoms coordinating with the manganese; and

Y is an oxidativaly-stable counterion.

- A composition according to claim 4, which comprises said paroxy compound at a laval of from 2 to 30% by weight and said catalyst at a leval corresponding to a manganase content of from 0.0005% to 0.5% by weight.
- A composition according to claim 4, wherein said manganese contant is from 0.001 to 0.25% by weight.
 - A composition according to claim 4, wherein said peroxy compound is salacted from the group consisting
 of hydrogen paroxide, hydrogen peroxide-liberating compounds, hydrogen peroxide-generating systems,
 paroxyacids and their salks, and peroxyacid bleach precursors, and mixtures thereof.
- A composition according to claim 4, which further comprises a surface-active material in an amount form 0.5% to 50% by weight.
 - A composition according to claim 4, which further comprises a datargancy builder in an amount from 5 to 80% by weight.
 - A composition according to claim 4, which furthar comprises an enzyma salacted from the group consisting of proteases, cellulases, lipases, amylases, oxidases and mixtures thereof.

11.	A method is provided for bleaching a stained substrate the method comprising contacting the stained sub-
	strate in an aqueous medium with a peroxy compound and a manganese complex each in an effective
	amount to interact with one another and provide a cleaning effect upon the substrate, the complex having
	the formula:

[LMn^{IV} (OR)₃] Y, (A)

wherein Mn is manganese in the +4 oxidation state;

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R is a C_1 - C_2 0 radical selected from alkyl, cycloalkyl, aryl, benzyl and radical combinations thereof, at least two R radicals may also be connected to one another so as to form a bridging unit between two oxygens that coordinate with the manganese;

- Lis a ligand selected from a C₂-C₆₀ radical having at least 3 nitrogen atoms coordinating with the manganese; and Y is an oxidatively-stable counterion.
- 12. A method according to claim 11, wherein the substrate is selected from fabric, dishware and dentures.
- 13. An aqueous medium comprising the manganese catalyst of claim 1 at a level from 0.001 ppm to 100 ppm of manganese and a peroxy compound that delivers active oxygen at a level from 0.01 ppm to 250 ppm.